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Deconvoluting the Complexity of Bone Metastatic Prostate Cancer via Computational Modeling

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The views, opinions and/or findings contained in this report are those of the author(s) and should not be construed as an official Department of the Army position, policy or decision unless so designated by other documentation.

# 1. INTRODUCTION:

TGF $\beta$  is a key regulator of bone development and is a key regulator in the vicious cycle of Prostate to Bone Metastasis. However, from a therapeutic standpoint, targeting TGF $\beta$ presents a dilemma given its noted pleiotropic roles. We hypothesize that an integrated computational modeling approach can be used to predict the temporal behavior of bone metastatic prostate cancer heterogeneous for TGF $\beta$  and further can be used to test the efficacy of putative therapies based on TGF $\beta$  inhibition. Multiple simulations and robust statistical post analyses will determine the role of TGF $\beta$  in the process. Our goal is to obtain a computational model of bone metastatic prostate cancer that can rapidly assess a putative TGF $\beta$  therapy in regards to efficacy and how to synergize with existing therapies to tackle heterogeneous metastases. This specialized research will be ultimately be used develop the PI professionally transition into an independent researcher.

# 2.KEYWORDS:

Provide a brief list of keywords (limit to 20 words). TGFβ- Transforming Growth Factor Beta Prostate Cancer (PCa) to Bone Metastasis Vicious Cycle Integrated computational modeling approach Heterogeneous metastases HLMCC- H. Lee Moffitt Cancer Center IMO- Integrated Mathematical Oncology pSMAD2- Phospho Smad2 staining technique

# 3. ACCOMPLISHMENTS:

Major goals and objectives of the project:

The major accomplishments were the prediction of the evolution of bone metastatic prostate cancers that are heterogeneous for TGF $\beta$  expression and utilization; and the exploration of the impact of a novel TGF $\beta$  inhibition on the progression of such heterogeneous bone metastatic prostate cancers. For this, the existing computational platform has been extended to consider TGF $\beta$  heterogeneity. Information derived from our collaborative studies with the Lynch Lab and the Moffitt Clinic were used to parameterize and calibrate the model.

Accomplishments under these goals:

35 human specimens of metastatic castrate resistant prostate cancer were obtained and examined. From this, quantification of the activity of pSMAD2 in the tissue revealed the differential expression of TGF $\beta$  ligands and the TGF $\beta$  receptors I and II (T $\beta$ RI and T $\beta$ RII), which were used by the PI to power the model and recapitulate the heterogeneity of the disease in the patient. Following this novel approach, the PI was able to explore different dosing strategies for a TGF $\beta$  inhibitor using the computational model of bone metastatic prostate cancer (Figure 1). As proposed in the original Statement of Work, the model was successfully used to inform the relevant biological pre-clinical mouse models of bone metastatic prostate cancer in the Lynch Lab. The PI published these results in Nature Scientific Reviews.



Figure 1. In silico effects of TGF $\beta$  inhibition on heterogeneous bone metastatic prostate cancer. (a) Individual human specimen of bone metastatic prostate cancer (Patient 41458) co-stained for TGF $\beta$  (green) and T $\beta$ RII (red). (b) Graph of staining pixel intensity for each clonal population. (c-f) In silico simulations (n $\geq$  24/group) were performed under control or TGF $\beta$  inhibition (80% efficacy) postand pre-treatment conditions. Clonal population was measured at Day 100 (left y-axis) and Day 250 (right y-axis).

The project provided the PI with the following opportunities for training professional development:

1) Structured mentoring committee program. Committee meetings were coordinated by the PI and held on a bi-monthly basis. These meetings assessed progress, scientific discussions of the challenges of the project, support on research activities and collaborations, resource availability and, manuscript and grant preparation. The mentoring committee comprised of Dr. David Basanta (Primary mentor), Dr. Alexander Anderson (Committee member) and Dr. Conor Lynch (Co-Mentor); experts with strong track record in the area of computational cancer modeling and cancer metastasis to bone collectively.

2) Environment to support the training plan. Through the IMO, the PI acquired the necessary soft/hardware to run simulations; and has used the cancer center's computer cluster and Biostatistics Core. In tandem, these allowed the PI to carry on the work described in the proposed research plan. Additionally, the co-Mentor, Dr. Lynch has provided invaluable assistance in the parameterization and validation of computational model outputs. The PI attended regular meetings with the genitourinary physicians (Drs. Julio Pow-Sang, Mayer Fishman and Shilpa Gupta) and pathologists (Dr. Jasreman Dhillon) and basic scientists (Drs. Nupam Mahajan and Jim Mulé).

*3) Education.* The HLMCC has provided the PI with feedback from an exceptionally active group of theoretical, clinical and basic faculty in cancer research. The PI will attended seminar series developed by the Cancer Biology and Evolution Program leaders, where he is learning modern modeling techniques and computational and biological tools for research from HLMCC faculty. A weekly Basic Science Grand Rounds and IMO seminars has also exposed the PI to international speakers, allowing the PI extensive opportunities to interact and learn from visiting scientists and mathematicians. Additionally the PI is attending relevant educational courses run by Oncological Sciences Department at the University of South Florida.

Results dissemination to communities of interest:

The PI has been responsible for the day-to-day management of the project and data collation. He has presented the generated data via committee meeting presentations, Lynch Lab weekly meetings, IMO departmental meetings, HLMCC Cancer Symposium short talks and HLMCC Post-Doctoral association meetings. The PI has maintained his membership to the American Association for Cancer Research (AACR), the Society for Basic Urology Research (SBUR) as well as to the Society of Mathematical Biology (SMB) and presented his data at the national meetings for these societies in either a poster or small talk format. The PI is

currently working on disseminate his research via social media and scientific blogging. The PI has attended HLMCC Post-Doctoral Association-organized grant and manuscript writing workshops. A research article has already emanated from the proposal, co-written with Cancer Biologist Dr. Leah Cook. The PI is currently writing a second manuscript and is preparing for his transition and search for an independent faculty position.

Plan for accomplishing the goals and objectives during the next reporting:

For the next reporting period, the PI is confident in generating a predictive computational model, not only go TGFB inhibition, but also its possible interaction with the current standard of care; as well as the dissemination of the basic biological and clinical understanding generated during this fellowship via conferences, workshops and social media. The PI also intends to begin his transition to an independent researcher, on schedule with that originally proposed.

# 4. IMPACT:

The development of the principal disciplines of the project:

The ability to rapidly assess the efficacy of the rapeutic strategies for incurable bone metastatic prostate cancer is an important and urgent need. Pre-clinical in vivo models provide insights yet are limited in their capacity to interrogate temporal multi-cellular interactions occurring in the cancer-bone microenvironment. In this project, the PI proposed that computational models that integrate current biological and clinical data can overcome this limitation and accelerate biological research. A direct result of this fellowship was a discrete continuum hybrid cellular automaton (HCA) model that can dissect the pleiotropic effects of inhibiting TGF<sup>β</sup> as a therapeutic target. Using our HCA model, we tested five different therapeutic doses delivered in two therapeutic windows. In silico results predict that TGF<sup>β</sup> inhibition, applied prior to tumor seeding acts by directly impacting prostate cancer cell viability but also by simultaneously restricting osteoclast formation and unexpectedly, promoting osteoblast differentiation. This effect was dependent on the prostate cancer cell expression of TGFB receptors. In silico predictions were validated with two independent in vivo models of bone metastatic prostate cancer (PAIII and C4-2B). Using immunohistochemical information from human bone metastatic prostate cancer samples, the PI alongside with the Lynch lab also demonstrated how the HCA can be used to predict the evolution of heterogeneous disease in response to applied therapies (Figure 1). Collectively, these data corroborate the overarching hypothesis of this work: that an integrated computational modeling approach can be used to predict the temporal behavior of bone metastatic prostate

cancer heterogeneous for TGF $\beta$  and further can be used to test the efficacy of putative therapies based on TGF $\beta$  inhibition.

Major Findings:

1. We have generated a novel computational model of bone metastatic prostate cancer that can be used to test and optimize new and extant treatments.

2. Using TGF $\beta$  inhibition as an example, we demonstrate how the computational model can be used to predict therapeutic windows in which inhibitors should be applied for maximum efficacy.

3. In control and treatment simulations, the computational model captures key temporal changes in the bone microenvironment and provides both high resolution qualitative and quantitative outputs.

4. The computational model describes for the first time the simultaneous pleiotropic effects of TGF $\beta$  inhibition on the cancer and bone compartment and reveals insights into TGF $\beta$  regulation of osteoblast/osteoclast function in the normal and bone metastatic microenvironment.

5. In vivo experimental findings with two independent models validate computational model outputs and also allow for refinement and improved accuracy of the model.

The computational model allows for the introduction of heterogeneous metastases (based on human tissue information) whose response to targeted therapies such as TGF $\beta$  inhibition can be analyzed. This has significant implications for patient-calibrated treatment strategies.

Impact on other disciplines:

The PI has developed new mathematical entities, of interest to the field of mathematics. During this work, a new agent-based model and analysis tools were developed, all offering new insights to the field of computer science and data science. Finally, the results have had impact on the clinic, as this work was highlighted in the last Annual Prostate Cancer Symposium; a meeting where clinicians, researchers and patients come together to assess the new discoveries and assess the next hurdles.

Technology transfer:

The PI is currently working on translating this new piece of research into an app that can be run on commercially available platforms.

Society beyond science and technology:

The PI has organized a Science Festival (Pint of Science Tampa), where this and other relevant themes were translated and presented in a friendly format to a lay audience.

The PI is also currently working on a video where the science behind this research is explained. We hope to release this video on youtube under a creative commons license. With this, the PI aims to improve public knowledge and attitudes toward this kind of interdisciplinary research, as well as highlight new discoveries on cancer that might have an impact of behavior, practices, decision-making and policies.

5. CHANGES/PROBLEMS:

Changes in approach and reasons for change: Nothing to Report.

Actual or anticipated problems or delays and actions or plans to resolve them: Nothing to Report.

Changes that have a significant impact on expenditures: Nothing to Report.

Significant changes in use or care of human subjects, vertebrate animals, biohazards, and/or select agents: Nothing to Report.

6. PRODUCTS:

Publications:

• Cook, L. M., Araujo, A., & Basanta, D. & Lynch, C. C. Predictive computational modeling to define effective treatment strategies for bone metastatic prostate cancer. Scientific Reports 6. 2016 doi:10.1038/srep29384. Published. Acknowledgement of federal support (yes)

• Araujo A & Bastanta D. Hybrid Discrete-Continuum Cellular Automaton (HCA) model of Prostate to Bone Metastasis. bioRxiv. 2016 Jan 1:043620. Published. Acknowledgement of federal support (yes)

Presentations:

• Society for Mathematical Biology Annual Meeting 2016, Nottingham, UK. "Predictive modeling of treatment strategies for bone metastatic prostate cancer"

• The 11<sup>th</sup> American Institute of Mathematical Sciences Conference on Dynamical Systems, Differential Equations and Applications 2016, Orlando, FL, USA. "Predictive modeling of treatment strategies for bone metastatic prostate cancer"

• Florida Prostate Cancer Research Symposium 2016, Clearwater, FL, USA. Predictive modeling of treatment strategies for bone metastatic prostate cancer"

Website(s) or other Internet site(s):

<u>arturoaraujo.com</u> - Repository of updates and main connection to the rest of social media distribution of material.

Technologies or techniques:

Agent-base Hybrid Discreet Cellular Automata- This technique is at the interface between mathematics and computer science, cross-linked by a visual platform written in java. This software was build by the PI and modified to receive relevant biological data and produce a clinically relevant simulation. The mathematical and computational framework was published in an open journal Biorxiv. The source code will be available on Github at the end of the project, when a stable final version is reached.

Inventions, patent applications, and/or licenses: Nothing to Report

Other products: Nothing to Report

# 7. PARTICIPANTS & OTHER COLLABORATING ORGANIZATIONS

Individuals that have worked on the project:

Name: Dr Arturo Araujo

Project Role: Principal Investigator

Researcher Identifier (e.g., ORCID ID): orcid.org/0000-0002-0269-2819

Nearest person month worked: 11 calendar months

Contribution to Project: Dr Araujo is an interdisciplinary scientist with a strong background in applying mathematical modeling to understanding cancer progression and joined Dr. Basanta's group in 2012. Dr. Araujo holds a PhD from University College London where he performed experimental and computational modeling of the role of aneuploidy in cancer. Dr. Araujo played a central role in the development of the initial Bone Modeling Unit model and his expertise in developing hybrid cellular automata has been critical in the development and understanding of the new model. He is the PI of the current proposal and will present his data at regular Basanta and Lynch lab meetings and report directly to his mentor Dr. Basanta

Funding Support: Department of Defense Prostate Cancer Research Program (W81XWH-15-1-0184)

# Name: Dr David Basanta

Project Role: Mentor

Researcher Identifier (e.g., ORCID ID): http://orcid.org/0000-0002-8527-0776 Nearest person month worked: 1 calendar month

Contribution to Project: Dr. Basanta is an Associate Member at the Moffitt Cancer Center and faculty in the Integrated Mathematical Oncology (IMO) Department, and currently serves as mentor for the PI. Dr. Basanta is a computer scientist with several years of experience in studying evolutionary dynamics in cancer. As a result of his role in the NCI's Integrative Cancer Biology Program (ICBP), Dr. Basanta is uniquely positioned to drive projects in which computational models, biological experiments and clinical parameters are combined and integrated with each other. Dr. Basanta is responsible for the overall management of the mathematical/computational aspects of the project. Dr Basanta is key in interpreting and integrating data from the Lynch lab. He works closely with the PI and Dr. Lynch and discusses plans and future directions with them on a regular basis. Dr. Basanta co-ordinates closely with the PI and Dr. Lynch in the dissemination of the collected data via publications in high impact journals. Funding Support: 1U01EB021664-01

# Name: Dr Conor C. Lynch

Project Role: Co-Mentor

Researcher Identifier (e.g., ORCID ID): orcid.org/0000-0002-4506-6244

Nearest person month worked: 1 calendar month

Contribution to Project: Dr. Lynch is an Associate Member in the Tumor Biology Department at the Moffitt Cancer Center since January 2011. Dr. Lynch currently serves as the co-mentor and provides all of the biological data proposed in this application. He directs the processing of human specimens for immunofluorescence analysis and the biological experiments examining the role of TGF $\beta$  in driving prostate cancer evolution in the bone microenvironment. Further, his lab has carried out the experiments informed by the computational model geared towards dissecting the role of TGF $\beta$  in prostate to bone metastasis. He has been responsible for the development of experimental strategies, interpretation of results, and the drafting of manuscripts. He has worked closely with the PI and Dr. Basanta in discussing his research strategy, results and future directions on a regular basis. For the past 3½ years, Dr. Lynch has directed a research program investigating the molecular mechanisms that facilitate prostate tumor-bone communication and has particularly focused on the role of matrix metalloproteinases (MMPs) in this process. He has extensive experience in animal husbandry, survival surgeries, animal care, animal imaging, DNA cloning, mutagenesis,

retroviral infection techniques, histological procedures and protein/DNA analysis. These techniques are integral to the proposed studies and the PI has a clear understanding of the techniques involved.

Funding Support: 1U01EB021664-01

Changes in the other active support of the PD/PI(s) or senior/key personnel since the last reporting period: Nothing to Report

Other organizations involved as partners: Nothing to Report

8. SPECIAL REPORTING REQUIREMENTS: None

9. APPENDICES:

# SCIENTIFIC REPORTS

Received: 19 April 2016 Accepted: 17 June 2016 Published: 14 July 2016

# **OPEN** Predictive computational modeling to define effective treatment strategies for bone metastatic prostate cancer

Leah M. Cook<sup>1,\*</sup>, Arturo Araujo<sup>2,\*</sup>, Julio M. Pow-Sang<sup>3</sup>, Mikalai M. Budzevich<sup>4</sup>, David Basanta<sup>2,+</sup> & Conor C. Lynch<sup>1,+</sup>

The ability to rapidly assess the efficacy of therapeutic strategies for incurable bone metastatic prostate cancer is an urgent need. Pre-clinical in vivo models are limited in their ability to define the temporal effects of therapies on simultaneous multicellular interactions in the cancer-bone microenvironment. Integrating biological and computational modeling approaches can overcome this limitation. Here, we generated a biologically driven discrete hybrid cellular automaton (HCA) model of bone metastatic prostate cancer to identify the optimal therapeutic window for putative targeted therapies. As proof of principle, we focused on TGF $\beta$  because of its known pleiotropic cellular effects. HCA simulations predict an optimal effect for TGF $\beta$  inhibition in a pre-metastatic setting with quantitative outputs indicating a significant impact on prostate cancer cell viability, osteoclast formation and osteoblast differentiation. In silico predictions were validated in vivo with models of bone metastatic prostate cancer (PAIII and C4-2B). Analysis of human bone metastatic prostate cancer specimens reveals heterogeneous cancer cell use of TGF $\beta$ . Patient specific information was seeded into the HCA model to predict the effect of TGF $\beta$  inhibitor treatment on disease evolution. Collectively, we demonstrate how an integrated computational/biological approach can rapidly optimize the efficacy of potential targeted therapies on bone metastatic prostate cancer.

Metastatic castrate resistant prostate cancer (mCRPC) typically manifests in the skeleton and is currently incurable<sup>1,2</sup>. In the bone microenvironment, prostate cancer cells hijack the normal bone remodeling process to create a "vicious cycle" of extensive bone formation and destruction<sup>3</sup>. Key mechanisms facilitating the cross-talk between the cancer and host compartment include the induction of receptor activator of nuclear  $\kappa B$  ligand (RANKL) expression and the release of sequestered growth factors from the bone matrix. Bone is a rich source of transforming growth factor $\beta$  (TGF $\beta$ ) and the role for this pleiotropic factor in promoting the survival and growth of bone metastatic cancers has been well described<sup>4,5</sup>. The molecular complexity of the circuitry driving this cycle has expanded tremendously in the past two decades revealing many potential targets for therapeutic intervention. The question remains however as to how to translate these potential therapies to the clinic. Biological experimentation and pre-clinical mouse models can be used to define the impact of putative therapies but are limited in their ability to dissect the potential dynamic and simultaneous effects on the multi-cellular tumor-bone microenvironment. One potential alternative approach is the integration of experimentally measured biological parameters with computational models to tackle the multi-scale nature of the disease<sup>6</sup>. Numerous computational models successfully demonstrate the feasibility of the approach<sup>7-14</sup>. Starting from existing experimental or clinical data it is possible to use statistical frameworks such as Approximate Bayesian Computation (ABC) to identify, in a "top-down" manner, the importance of unknown parameters in disease progression by applying a distribution of probability on those factors<sup>15</sup>. Conversely, agent based models, such as discrete-continuum Hybrid Cellular

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Automata (HCA), are better suited to test hypotheses using a mechanistic "bottom-up" approach to provide unbiased predictions<sup>16</sup>. These models work by parameterizing the properties of cells (or agents) with regards to proliferation, apoptosis, secretion of factors, genetic mutations or even metabolism<sup>17</sup>. The ability to apply HCA models to two- or three-dimensional grids make them uniquely qualified for studying temporal tumor-host interactions over time, especially in the context of applied therapies<sup>15,18-20</sup>.

Previously, our group generated a HCA based computational model of the bone modeling unit (BMU) that recapitulates the homeostatic sequence of bone resorption and anabolism<sup>18</sup>. The *in silico* BMU is 1000  $\mu$ m × 1500  $\mu$ m and is composed of bone, mesenchymal stromal cells (MSCs), precursor and adult osteoblasts, and precursor and mature multinucleated osteoclasts. The sequence and timing of resorption and bone formation that emerges from the model recapitulates the extensive literature and the interactions of the cells were carefully modeled around bone derived factors including RANKL and TGF $\beta^{18}$ . Using human parameters based on the growth of prostate cancer in bone we demonstrated that the introduction of an emboli of prostate cancer cells ( $\geq$ 9) into the BMU was sufficient to consistently initiate the vicious cycle. Subsequently, cancer-bone interaction could be monitored over a clinically relevant 250-day period<sup>21</sup>. We also reported how the model could be used to potentially optimize the effects of bisphosphonates and anti-RANKL therapies that are components of the current standard of care. In the current study, a major objective was to use the model to explore the impact/ efficacy of putative inhibitors. Our previously published HCA model, as expected, defined an important role for TGF $\beta$  in regulating cancer-bone interaction<sup>18</sup>.

TGF $\beta$  inhibitors such as neutralizing antibodies are currently undergoing clinical trial<sup>22</sup>. However, their application for the treatment of osteogenic bone metastatic prostate cancer has not been explored thus far due to the pleiotropic and often opposing effects TGF $\beta$  can have on cancer and bone cell behavior<sup>5,23–25</sup>. Therefore, we posit that TGF $\beta$  inhibition would be an ideal challenge for testing the predictive power of our HCA based model. Here, using an evolved version of the HCA model, we simulated various therapeutic strategies (i.e. inhibitor concentration, time of therapeutic intervention) to predict the optimal efficacy of TGF $\beta$  inhibition. Further, the enhanced HCA model provides new insights into how TGF $\beta$  can regulate multi-cellular interactions over time. HCA outputs were validated *in vivo* using two models of osteogenic bone metastatic prostate cancer. Moreover, using patient specific information from bone metastatic specimens, we demonstrate the flexibility of the HCA model in predicting the efficacy of TGF $\beta$  inhibitors on lesions that are heterogeneous for TGF $\beta$  utilization. Collectively, we demonstrate how an integrated computational/biological modeling approach can be used to optimize therapy efficacy for the treatment of bone metastatic prostate cancer.

#### Results

Computational modeling of TGFB inhibition in normal bone remodeling and in bone metastatic **prostate cancer.** TGF $\beta$  is known to have concentration dependent pleiotropic effects on osteoblasts and osteoclasts<sup>26-28</sup>. In silico, the ability of stromal cells to respond to varying TGF<sup>β</sup> concentrations (0.1 to 10 ng/ml) was integrated into our HCA of normal bone remodeling, the bone multicellular unit (BMU) (see Equations 1, 2, and Table S1)<sup>18</sup>. To obtain a realistic readout of the level of TGF $\beta$  inhibition that could be achievable *in vivo*, we treated mice with a TGF $\beta$  neutralizing antibody (1D11) at a dose previously used in the literature (10 mg/kg) and consistent with clinical trials performed for a humanized version of the 1D11 TGF $\beta$  neutralizing antibody, fresolimumab/GC1008<sup>22,29,30</sup>. We observed that the TGF $\beta$  neutralizing antibody significantly reduced circulating TGF $\beta$  serum levels by up to 80% compared to IgG control treated animals (Supplementary Fig. S1). Using phospho-SMAD2 as a surrogate for TGF $\beta$  activity<sup>31</sup>, we also observed that the TGF $\beta$  neutralizing antibody could inhibit TGF $\beta$  activity 50–80% in tumor naïve and tumor bearing tissues (Supplementary Fig. S1 and Fig. 3f). Based on this *in vivo* information, we applied the TGF $\beta$  inhibitor to the normal BMU at a level of 80% *in silico*. The stochastic nature of the BMU allows for variation and statistical analysis of simulation outputs. The results of multiple simulations (n = 29/group) show that TGF $\beta$  inhibition significantly promoted bone formation (9% increase) over a 75-day period by enhancing osteoblast expansion and differentiation while limiting osteoclast viability (Fig. 1a, Supplementary Fig. S1, and Table S2). Importantly, these in silico results are consistent with previous *in vivo* studies and support the robustness of the parameters used to power the computational model<sup>29,32</sup>.

The vicious cycle paradigm suggests that metastatic prostate cancer cells utilize TGF $\beta$  signaling to promote their survival and growth. We therefore seeded the computational model with TGF $\beta$  responsive prostate cancer cells. Once the vicious cycle was established at day 80, we initiated TGF $\beta$  inhibition (post-treatment scenario). Simulations (n = 24/group) revealed that TGF $\beta$  inhibition reduced cancer growth by approximately 15%, but only when the inhibitor was applied at a constant 99% level of efficacy until day 250 (Supplementary Fig. S2 and Video 1). At a more biologically relevant level of 80% inhibition, we observed little difference in cancer cell growth between the control and treatment groups. Surprisingly, and in contrast to the observed effects of TGF $\beta$  inhibition on the normal BMU, we also observed no difference in osteogenesis between the control and treatment groups even at later stages (Supplementary Fig. S2, Table S3, and Video 1). Taken together, these results suggest that the treatment of established and actively growing bone metastases with TGF $\beta$  inhibitors would have no impact on the progression of the lesions unless the inhibitor was >99% effective.

A major advantage of the computational model is that it can be used to explore therapeutic windows of efficacy for putative inhibitors. Simulations (n = 24) revealed that applying the TGF $\beta$  inhibitor *in silico* at day 1 prior to the seeding of the cancer cells (pre-treatment scenario) even at a level of 20% efficacy significantly reduced tumor burden over time by  $\geq$ 65% (Fig 1b, Supplementary Fig. S2, Table S3, and Video 2). Interestingly, TGF $\beta$  inhibition resulted in a small but significant increase in cancer-induced osteogenesis compared to control during early tumor progression (Day 100). However, at the end of the simulations there was significantly less cancer-induced osteogenesis in the TGF $\beta$  treated group compared to control (Fig. 1b, Supplementary Fig. S2, and Table S3, Day 250).



**Figure 1.** *In silico* effects of TGF $\beta$  inhibition on normal and prostate cancer induced bone turnover (**a**,**b**) *In silico* control and TGF $\beta$  inhibitor treated simulations in normal (BMU, n = 29/group, **a**) and bone metastatic prostate cancer (PCA-Bone Mets, n = 24/group, **b**) scenarios. Representative images of simulation runs at indicated time points are shown with magnified insets. TGF $\beta$  inhibitor was applied at day 1 for all simulations (pre-treatment scenario). Cell populations analyzed include mesenchymal stem cell (MSC), osteoblast precursors (pOB), osteoblasts, osteoclast precursors (pOC) and osteoclasts. Temporal changes in bone area ( $\mu$ m<sup>2</sup>) were also predicted under control and TGF $\beta$  inhibitor conditions.

These *in silico* results predict that applying TGF $\beta$  inhibitors in a preventative manner will reduce the growth of metastatic prostate cancer without exacerbating cancer induced osteogenesis.

*In vivo* validation of computational model predictions. Analysis of human specimens of bone metastatic prostate cancer derived from deidentified cancer patients at the Moffitt Cancer Center (n = 20) show that TGF $\beta$  ligand and receptors are expressed and pSMAD2 staining indicates that the TGF $\beta$  pathway is active (Fig. 2a). *In vitro* analysis of prostate cancer cell lines that can grow in the bone microenvironment identified the PAIII cell line as reflecting the TGF $\beta$  receptor and growth factor producing (TRP) status observed in human specimens (Fig. 2b,c). We also noted that the PAIII cell line was sensitive to inhibition with TGF $\beta$  inhibitors (Fig. 2d,e). We therefore initially chose this cell line to test computational model predictions.

Dissecting cancer cell behavior *in silico* illustrates TGF $\beta$  inhibition directly limits growth over time by impacting cancer cell proliferation (Fig. 3a–d). To determine the validity of the computational outputs, 6-week-old male SCID Beige mice were pre-treated with either a TGF $\beta$  inhibitor (TGF $\beta$ i-1D11, 10 mg/Kg, 3× weekly; n = 10) or an isotype control IgG (Control-13C4, 10 mg/Kg 3× weekly; n = 8) and subsequently inoculated with luciferase-expressing PAIII cells. Bioluminescence analysis revealed a significant reduction in tumor growth in the TGF $\beta$  inhibitor treated group compared to controls and, as expected, reduced pSMAD2 and AKT phosphorylation (Fig. 3e,f). We further found significant reductions in proliferation (40%) and increases in apoptosis (70%) between the TGF $\beta$  inhibitor treated and control groups (Fig. 3g). The HCA model is based on humanized



Figure 2. TGF $\beta$  expression and utilization in prostate cancer specimens and cell lines. (a) Immunofluorescence of TGF $\beta$ , T $\beta$ RII, and pSMAD2 (red) in human (cytokeratin-green) bone metastatic prostate cancer (n = 20). Dashed box represents area of magnification. Graphs represent intensity of pixels. Scale bars represent 100 µm. (b,c) Real time PCR analysis of T $\beta$ RI and T $\beta$ RII expression (b) and ELISA measurement of TGF $\beta$  concentration (c) in PAIII, C42B and PC3. (d) The effect of increasing concentrations of TGF $\beta$  inhibitor (TGF $\beta$ i; 1D11 antibody) on SMAD reporter activity (RLU). (e) The effect of TGF $\beta$  inhibition (TGF $\beta$ i; 1D11 10 µg/ml) on colony formation and size compared to control (Control-13C4, 10 µg/ml). Asterisks denote statistical significance (\*p < 0.05; \*\*p < 0.01).

parameters and therefore, to compare computational model outputs to those obtained with the PAIII model, we used a scaling approach. Briefly, using tumor growth rates from the computational model (25 day intervals) and the animal model (2 day intervals) we calculated the slope of the line/derivative for each time point. Using this approach, we found that derivatives for each model are similar for pre and post treatment between day 8–10 for the animal model and day 75–100 for the *in silico* model. At these comparable points, the computational model accurately predicts the impact of TGF $\beta$  inhibition on cancer cell proliferation but differences in apoptosis were not evident in the computational model until later time points (Fig. 3d vs. 3g).

To further assess the predictive power of the computational model, we performed a TGF $\beta$  inhibitor post-treatment study. *In silico* findings suggest treatment of an established TGF $\beta$  ligand and receptor-expressing cancer (TRP) with a TGF $\beta$  inhibitor would not be of benefit. Our *in vivo* data confirm these predictions (Supplementary Fig. S3).

**TGF**β **inhibition prevents prostate cancer induced osteolysis.** Because of their role in the vicious cycle, we next focused on osteoclast behavior over time in the control and TGFβ inhibitor treated groups. Dissection of the computational model outputs revealed the number of active bone resorbing osteoclasts were significantly lower in the TGFβ inhibitor versus control group at day 100 (Fig. 4a). Further, the rate of osteoclast maturation and apoptosis was significantly mitigated during this period, which is in agreement with known effects of TGFβ inhibition on osteoclast function (Fig. 4b,c)<sup>28,33</sup>. Analysis of the lytic component of our *in vivo* model identified that there was approximately 50% (p=0.002) less tumor induced osteolysis in the TGFβ inhibitor group compared to the controls as determined by X-ray (Fig. 4d). Histochemical analysis using the osteoclast specific marker tartrate resistant acid phosphatase (TRAP) demonstrated that this reduction in osteolysis was due to significantly fewer osteoclasts in the TGFβ inhibitor treated group compared to the controls (Fig. 4e). These data suggest TGFβ inhibition reduces the extent of cancer-induced osteolysis over time *in vivo* by limiting osteoclast function and validates the pre-treatment HCA model outputs.

**TGF** $\beta$  inhibition has differential effects on normal and cancer induced bone formation. Based on published data demonstrating that TGF $\beta$  blockade increases bone formation<sup>29</sup>, we hypothesized that inhibition of the growth factor would significantly exacerbate prostate cancer induced osteogenesis. At day 100, the computational model does predict a small but significant increase in bone formation in the TGF $\beta$  inhibitor simulations despite a concomitant decrease in osteoblast proliferation (Fig. 5a–c and Supplementary Fig. S2). While



Figure 3. TGF $\beta$ i pre-treatment prevents bone metastatic prostate cancer growth *in silico* and *in vivo*. (a) Representative *in silico* outputs from control (left panel) and TGF $\beta$ i (right panel) treated simulations at day 100. (**b**-**d**) *In silico* predictions of TGF $\beta$  inhibition on cancer cell growth (**b**), and cancer cell proliferative/apoptotic rates (average number of proliferating/apoptotic cancer cells at 25 day intervals over a 250 day period, (**c**,**d**). (**e**) Bioluminescence measurement of PAIII growth under TGF $\beta$ i (1D11, 10µg/ml; n = 10) or control (13C4, 10µg/ml; n = 8) conditions. (**f**) pSMAD2 and pAKT positivity (red) as a ratio of unphosphorylated protein. Scale bars, 25 µm. (**g**) The proliferative and apoptotic index in TGF $\beta$ i and control tissue sections were measured using pHistone H3 and cleaved caspase-3 (c-Caspase-3) (red) respectively as a ratio to total cell number (DAPI; blue). Scale bars, 50 µm. Asterisks denote statistical significance (\*p < 0.05; \*\*p < 0.005).

microCT ( $\mu$ CT) scans of tumor bearing bones did not reveal any differences between the groups (Fig. 5d), histomorphometry analysis does support the *in silico* predictions with increased bone formation in TGF $\beta$  inhibitor treated mice and a trend towards fewer bone rimming osteoblasts (Fig. 5e,f). In contrast, we observed a robust increase in bone formation in contralateral sham limbs derived from the TGF $\beta$  inhibitor group for all measurements (Fig. 5d–f). Taken together, these *in vivo* data suggest that TGF $\beta$  inhibition does not greatly exacerbate prostate cancer induced osteogenesis compared to non pathological conditions and support, in part, HCA model outputs. It is also worth noting that over a longer period (Day 250), the computational model predicts TGF $\beta$  inhibition would ultimately result in decreased bone formation (Fig. 5a and Supplementary Fig. S2), a finding that warrants further exploration *in vivo*.

**Predicting TGF** $\beta$  inhibitor efficacy on heterogeneous bone metastatic prostate cancer. While the majority of cancer cells in human specimens of bone metastatic prostate cancer produce both TGF $\beta$  ligand and receptors (TRP), we also noted the presence of cancer cells that either produced the ligand alone (TP), the receptor alone (TR) or neither (TN) (Fig. 2). This raises the question as to what the impact of TGF $\beta$  inhibition would be on these clonal populations. We have shown that, in our hands, the C4-2B cell line produces TGF $\beta$  but not the receptors (TP) (Fig. 2b–d). Therefore, we challenged the computational model to determine the impact of TGF $\beta$  inhibition on a homogenous TP bone metastatic cancer and found no effect on the growth of the cancer in pre- or-post treatment simulations (Equation 5, Supplementary Fig. S4). To test these results *in vivo*, C4-2B luciferase-expressing cells were inoculated into mice that were pre- or post-treated with TGF $\beta$  inhibitor or IgG control and we observed no difference in tumor growth between the groups (Supplementary Fig. S4).

Clinically, human samples of bone metastatic prostate cancer are heterogeneous for their usage of  $TGF\beta$  (Fig. 2a). A major advantage of the HCA modeling approach is the ability to integrate multiple clonal phenotypes



**Figure 4. TGF** $\beta$ **i effects on cancer induced osteolysis. (a)** Osteoclast activity in the TGF $\beta$ **i** and control groups was determined *in silico* at 25 day intervals over a 250 day period. (**b**,**c**) The effect of TGF $\beta$ **i** versus control on osteoclast fusion/maturation and apoptosis over time as a ratio to total cell number. (**d**) Ratio of tumor volume (measured by total area of osteolysis; TuV) to total volume (TV) in X-rays of tumor and sham tibia from TGF $\beta$ **i** (n = 10) and control (n = 8) groups. (**e**) Quantitation of TRAcP (red) positive osteoclasts per tumor/bone interface. Asterisks denote statistical significance (\*p < 0.05; \*\*p < 0.005; \*\*\*p < 0.0001).

based on patient specific information. In this regard, analysis of a single patient specimen, identified heterogeneous expression and activity in the TGF $\beta$  signaling axis with the following clonal ratio noted: TP 1: TRP 231: TR 6: TN 4 (Fig. 6a,b). We seeded the HCA model with this ratio of clones and performed simulations for control, pre- and post-treatment conditions (n  $\geq$  24/group) to determine the impact of TGF $\beta$  inhibition on metastatic prostate cancer evolution (Equations 3–6). *In silico* data show the effects of pre- and post- TGF $\beta$  inhibitor treatment on clonal evolution for this patient over time and the responses of the surrounding tumor microenvironment (Supplementary Fig. S5, Table S4, and Video 3). The model also predicts the dominance of the TRP clone over time under control conditions (Fig. 6c). However, application of the TGF $\beta$  inhibitor, especially in the pre-treatment groups at varying levels of efficacy, show how the cancer cell dynamics shift over time in favor of other clones, in particular for the TN population (Fig. 6d–f and Table S4). These data underscore the power of the HCA computational modeling approach in examining the temporal effects of targeted therapy on heterogeneous cancer cells and the surrounding microenvironment.

#### Discussion

Current treatment options for patients diagnosed with bone metastatic castrate resistant prostate cancer include second generation androgen deprivation therapies, radiation treatment, bisphosphonates/anti-RANKL therapies, alpharadin and/or chemotherapy<sup>1</sup>. While these treatments mitigate pain, pathological fracture and increase overall survival, the disease remains incurable with the median survival time being approximately 3-years subsequent to diagnosis. Increasing our knowledge of the mechanisms driving the disease can reveal novel therapeutic targets. To this end, dozens of molecular mechanisms that play important roles in tumor-bone interaction have been discovered<sup>34</sup>. However, dissecting how potential targeted therapies will work in the context of current treatment paradigms and their translation to the clinical setting presents a major challenge. Using TGF $\beta$  inhibition as an example, we demonstrated how a novel biologically driven computational HCA model can rapidly define temporal cancer-bone microenvironment responses to a given therapy. Further, the integrated approach provides insight into optimal therapeutic windows to apply a given inhibitor. For TGF $\beta$  inhibition, the HCA model indicates that application of the inhibitor in an adjuvant setting subsequent to the detection/treatment of aggressive prostate cancer would be most effective.

As with biological models, there are numerous mathematical approaches to study cancer progression such as branching and Moran processes based models, systems of ordinary and partial differential equations, and agent based models, each with varying strengths and weaknesses<sup>6,15,35,36</sup>. Data driven "top down" models "fit" existing clinical or experimental information to identify parameters that explain the behavior of the disease<sup>37,38</sup>. In the context of prostate cancer, many elegant models have used biological parameters such as prostate specific antigen (PSA) to predict time to progression, and to model the effects of intermittent androgen therapy<sup>8</sup>. The tightly regulated process of normal bone remodeling lends itself well to modeling how key factors such as RANKL and TGF $\beta$  control the behavior and activity of bone stromal cells over time<sup>39–43</sup>. By extension, perturbing this balanced ecosystem with an invasive species such as cancer can also be modeled. Our own group, as well as others, has been exploring how bone metastatic cells and skeletal malignancies such as myeloma interact with the bone microenvironment in order to progress<sup>18,44,45</sup>. In the current study, we used an agent-based approach that allows



Figure 5. TGF $\beta$ i impact on osteogenesis. (a) *In silico* changes in bone area ( $\mu$ m<sup>2</sup>) were predicted at 25-day intervals over a 250 day period. (b,c) The effect of TGF $\beta$  inhibition on the average number of proliferative and apoptotic osteoblasts per day *in silico*. (d)  $\mu$ CT analysis of bone volume (arrow) to total volume (BV:TV) in control and TGF $\beta$  inhibitor treated bones. (e,f) Histomorphometry analysis of osteogenesis (e) and number of bone rimming osteoblasts (arrows; f). Asterisks denote statistical significance (\*p < 0.05; \*\*p < 0.005; \*\*\*p < 0.0001).

for the exploration of key cellular interactions over space and time in an unbiased hypothesis driven "bottom-up" manner. We have shown how experimentally-derived cellular parameters can be integrated into the rule sets and partial differential equations (PDEs) used to drive HCA models. This allows us to produce biologically<sup>46</sup> and clinically<sup>47</sup> testable hypotheses without making assumptions regarding population level dynamics. The result is a model capable of generating predictions that naturally emerge from the interactions between cells and their environment. Key to the robustness of the HCA model outputs, is the reliability of the biological parameters used to power the PDEs. In the current study, we used human parameters to generate the HCA model<sup>21</sup>. While we assume that individual bone stromal cell components have similar dimensions and lifespans between the species, human cancer cells typically grow much more slowly than human derived xenografts or animal models of the disease<sup>21</sup>. Therefore appropriate scaling between the models is an important consideration for the comparison of results. Further, not all of the HCA models predictions were correct. For example, we observed the *in silico* effects of TGF $\beta$  inhibition on osteoblast number was discordant with *in vivo* results (Fig. 5b vs. 5f), suggesting that the parameters/assumptions governing the effects of TGF $\beta$  inhibition on apoptosis require re-calibration based on the obtained biological data. This reiterative process allows for fine-tuning of the HCA.

The roles of TGF $\beta$  in skeletal development and malignancy have been well described<sup>5,24</sup>. However, enthusiasm for applying TGF $\beta$  inhibitors as a therapeutic strategy to treat metastatic bone disease including mCRPC is limited because of the pleiotropic and often opposing roles TGF\beta plays in normal and cancer cell biology<sup>5,24,25,48,49</sup>. This complexity made TGF $\beta$  and the effects of TGF $\beta$  inhibition an ideal challenge with which to test the power of our HCA based computational model. Our results indicate the treatment of established active metastases with a TGF<sup>β</sup> inhibitor, unless applied at >99% efficacy, would have little or no impact on the progression of cancer cells regardless of their dependency on TGF $\beta$ . In contrast, the application of the inhibitor in a preventative or adjuvant manner would significantly control bone metastatic prostate cancer growth and osteogenesis. This result was predicated on the basis that the metastatic cancer cells have an active TGF $\beta$  signaling axis which we found to be the case in the majority of human bone metastatic prostate cancer specimens. The mathematical model was built on the assumption that  $TGF\beta$ , RANKL and bone derived nutrients drive cell responses. Arguably then, interfering with TGFB signaling in the model could potentially be self-fulfilling in predicting cancer-host behavior. However, despite the limited number of cytokines and growth factors included in the model, we validated many of the model predictions with independent *in vivo* experiments thus reinforcing the key roles for TGF $\beta$  and RANKL in the vicious cycle of tumor bone interaction. The majority of our in vivo results support the accuracy of the parameters and assumptions used to power the computational model. The in vivo results are consistent with other studies examining the role of TGF $\beta$  in bone metastatic cancers including prostate, breast and melanoma but, importantly, underscores the potential for computational modeling in predicting the efficacy of an





applied targeted therapy<sup>50–52</sup>. Although our model is centered on the roles of TGF $\beta$  and RANKL, it is important to note that the circuitry of the HCA model can easily be expanded to include other cell types (immune cells or cancer-associated fibroblasts) and molecules (e.g. PTHrP, BMPs) and their roles/effects subsequently explored.

While the computational model was primarily used to examine TGF $\beta$  inhibition in bone metastatic prostate cancer, the outputs generated also revealed new insights into TGF $\beta$  biology. Firstly, the model highlights the critical role for TGF $\beta$  in the cyclical dynamics of populations such as osteoclasts and confirms the importance of the osteoclast in initializing the vicious cycle. For example, we noted osteoclast infiltration often precedes a period of cancer growth and osteogenesis (Video S2), a finding that could be informative for the timing of anti-resorptive therapies as we have previously shown<sup>18</sup>. TGF $\beta$  inhibition alters these dynamics and at specific time-points can reverse the trends between the treated and control groups so that osteoclast numbers in the control simulations may in fact be lower than those in the TGF $\beta$  inhibitor treated group (Fig. 3A). This suggests that arbitrarily selected time-points in pre-clinical *in vivo* animal studies may not accurately reflect how applied therapeutics are impacting cancer-bone interaction over time. Secondly, although TGF $\beta$  inhibition promotes robust bone formation in normal non-pathological situations, the model and our *in vivo* results confirm counter-intuitively that, TGF $\beta$  inhibition does not greatly exacerbate prostate cancer induced osteogenesis.

Currently, mCRPC inter-patient heterogeneity is a major clinical challenge. Integrating biological and computational modeling offers a unique opportunity to study how cancer evolves and reacts to changing microenvironments and applied targeted therapies. Patient derived xenograft (PDX) samples are being used to design precision treatment strategies and integrating the biological parameters derived from these specimens into computational models could prove to be a synergistic way to tackle the complexity of heterogeneity in individual patients<sup>19</sup>. Again, using patient derived TGF $\beta$  signaling axis information as an example, we demonstrated how clonal variation and evolution in response to applied TGF $\beta$  inhibitors could be incorporated into the HCA model (Fig. 6). In these studies, we assumed the TGF $\beta$  dependent growth rates of clonal cancer cells. For personalization of the HCA and its clinical application, individual patient specimens would have to be isolated and growth rates of various clones examined in *ex vivo* assays. Current advances in single cell *ex vivo* analyses support the feasibility of such an approach<sup>53,54</sup>. The HCA model also allows for the optimization of inhibitor dosing and timing that in turn could be used to generate an adaptive therapy strategy to prevent the outgrowth of resistant sub-populations<sup>20,55</sup>.

In conclusion, we have developed a novel and unbiased computational HCA model that allows for the dynamic multi-scale understanding of how metastatic prostate cancer cells evolve and interact with the surrounding bone microenvironment. We used this model to predict the efficacy and response of bone metastatic prostate cancer to targeted therapies such as, TGF $\beta$  inhibitors. Further, our integrated computational and biological approach allowed for the dissection of how TGF $\beta$  inhibition simultaneously affects osteoblast, osteoclast and cancer cell behavior over time. The HCA model constitutes a platform for discovery that can readily be expanded to incorporate additional cellular and molecular circuitry. This will ultimately yield a clinical tool that will aid the medical oncologist in designing curative strategies for heterogeneous bone metastatic prostate cancer. Most

importantly, this modeling approach can be applied to the development of new therapeutic strategies across a broad spectrum of human malignancies.

#### Methods

An HCA Model of bone metastatic prostate cancer for therapy optimization. In the computational model, each cell type responds to TGF $\beta$  levels in an either directly proportional  $(1 + Log(TGF\beta))$  or inversely proportional  $(-1Log(TGF\beta))$  manner. TGF $\beta$  inhibition in the HCA is achieved by controlling TGF $\beta$ bioavailability (0–99% inhibition at a constant level). Computational models were seeded with homogeneous or heterogeneous prostate cancer cells that expressed the TGF $\beta$  receptor and ligand (TRP), the ligand alone (TP), the receptor alone (TR) or were negative for both (TN). For the HCA model, we consider cellular intrinsic behaviors and the impact of TGF $\beta$  on these behaviors. We include 8 different cell types: resident and active cells of the bone stroma (mesenchymal stromal cells (MSCs), precursor osteoblasts (pOB), adult osteoblasts (aOB), precursor osteoclasts (pOC), adult osteoclasts (aOC)) and prostate cancer cells of varying TGF $\beta$  responsiveness (TRP, TGF $\beta$ ligand and receptor-producing; TR, receptor-producing; TP, ligand-producing and; TN, negative for receptor and ligand expression). We considered interactions between all cell types and the impact of these interactions on the tumor-bone microenvironment. Empirical, experimental and theoretical parameters were used to fuel equations (Table S1).

*Precursor Osteoblasts and Osteoclasts.* Based on empirical data and literature, we assume a maximum rate of pOB division as 36 hours based on ATCC specifications for MC3T3-E1 cells. The rate of pOB division is inversely affected by TGF $\beta^{28}$  and we assume that the effect is logarithmic. If TGF $\beta$  is at saturation levels (>10 ng/ml), then the pOB division rate tends to zero. This assumption is based on our findings of the effect of TGF $\beta$  on osteoblast precursors (Supplemental Fig. S1). If there is no TGF $\beta$  present, then the maximum rate is considered. The rate of division has subsequent effects on the number of mature bone generating osteoblasts. By the same token, the fusion rate of pOC is also affected inversely proportional to the availability of TGF $\beta$ . These behaviors can be described by:

$$p(precursor \ action) = Div_{precursor} \ (1 + 1/(0.1 * Log(TGF\beta)))$$
(1)

where  $0 < TGF\beta < 1$  and  $Div_{precursor}$  is substituted in the case of precursor osteoblasts,  $Div_{pOB}$  is the maximum rate of pOB division, and in the case of precursor osteoclasts,  $Div_{pOC}$  the maximum rate of pOC fusion in the absence of TGF $\beta$ . This ensures that, when bone is being resorbed and TGF $\beta$  is being made bioavailable, osteoclastogenesis is limited.

Once fused, the probability of aOC survival also depends proportionally on TGF $\beta$ . This is calculated by:

$$p(aOC \ survival) = Surv_{aOC}(-1/(0.1 * Log(TGF\beta)))$$
<sup>(2)</sup>

where  $0 < TGF\beta < 1$  and  $Surv_{aOC}$  is the maximum percentage of survival for aOCs (100% when TGF $\beta$  is as its saturation level). If the levels of TGF $\beta$  are below the saturation level, the probability of death for the aOC increases<sup>56,57</sup>.

Bone metastatic prostate cancer cells. Prostate cancer cells were explicitly defined as being dependent on TGF $\beta$  and bone derived nutrients (*BDN*) for their division. The probability of division was estimated as being proportional to the inverse logarithm of the available BDN. If there are no nutrients, there is zero division. If there is maximum nutrient saturation, the division rate is at its maximum.

Based on empirical data obtained with TRP cells such as PAIII, we assume that TRP cells have a maximum division rate of once every 1.5 days and a lifespan without contact with BDN of 14 days based on low serum (2%) soft agar assays (Supplementary Fig. S4). TRP division in response to bone derived nutrients was modeled as follows:

$$p(TRP \ division) = Div_{TRP}(1 + 1/(0.1 * Log(BDN)))$$
(3)

where  $0 < TGF\beta < 1$  and  $Div_{TRP}$  is the maximum rate of TRP division, in the saturation level of bone derived nutrients. Under TGF $\beta$  inhibition, the maximum division rate is assumed to be reduced for TRP to once every 2 days based on empirical observations in our laboratory (data not shown).

We assume that TR has a maximum division rate of 1.75 days, but a lifespan without nutrients of 10 days, calculated as not having a cost for producing TGF $\beta$  ligand but benefiting from the presence TGF $\beta$  in the bone microenvironment. TR cell division depends directly on bone-derived nutrients in the same manner:

$$p(TR division) = Div_{TR}(1 + 1/(0.1 * Log(BDN)))$$
(4)

where 0 < BDN < 1 and  $Div_{TR}$  is the maximum rate of TR division, in the saturation level of bone derived nutrients. Under TGF $\beta$  inhibition, the maximum division rate is reduced for TR to once every two days.

We assume that TP has a maximum division rate of once 1.75 days, and a lifespan without nutrients of 10 days, calculated as having a cost for producing TGF $\beta$  ligand but not benefiting from TGF $\beta$  contained within the bone derived nutrients. TP cell division depends directly on bone-derived nutrients in the same manner:

$$p(TP \ division) = (Div_{TP}1 + 1/(0.1 * Log(BDN)))$$
(5)

where 0 < BDN < 1 and  $Div_{TP}$  is the maximum rate of TP division. Under TGF $\beta$  inhibition, the maximum division rate of TP remains the same.

Finally, we assume TN has a maximum division rate of once every 2 days and a lifespan without nutrients of 12 days, calculated from having no cost of producing TGF $\beta$  and no benefit from TGF $\beta$  signaling. TN cell division depends directly on bone-derived nutrients through:

$$p(TN \ division) = (Div_{TN}1 + 1/(0.1 * Log(BDN)))$$
(6)

where 0 < BDN < 1 and  $Div_{TN}$  is the maximum rate of TN division. Under TGF $\beta$  inhibition, the maximum division rate of TN remains the same.

**Cell Culture and Patient Specimens.** Luciferase-expressing PAIII, C42B, and PC3 prostate cancer cell lines were cultured in complete Dulbecco's Modified medium supplemented with 10% fetal bovine serum<sup>58-60</sup>. All cell lines were periodically tested for mycoplasma (#CUL001B, R&D Systems) and short tandem repeat (STR) verified at the Moffitt Clinical Translational Research Core. De-identified tissue sections of bone metastatic prostate cancer were obtained from the Moffitt tissue archives (MCC 50086).

*In vivo* experiments. All animal experiments were performed with IACUC approval (R1283) and were conducted in accordance with the guidelines set forth in the *Guidelines for the Care and Use of Laboratory Animals* published by the National Institutes of Health. Pre-Treatment Studies: 6-week old male SCID Beige mice were injected intraperitoneally with either TGF $\beta$  inhibitor (1D11; 10µg/ml; n = 10/group) or isotype control, 13C4 (10µg/ml; n = 8/group), a kind gift from Scott Lonning and Patrick Finn at Genzyme. Subsequently, luciferase expressing PAIII or C4-2B cell lines (5 × 10<sup>4</sup> or 1 × 10<sup>5</sup> respectively in 20µl of saline) were intratibially injected either one-day or one week after TGF $\beta$  pre-treatment<sup>61,62</sup>. Contra-lateral limbs were injected with saline and served as a positive control. Mice received TGF $\beta$  inhibitor or IgG control injections every three days (PAIII model) or weekly (C4-2B). Bioluminescence was measured longitudinally as a correlate of tumor growth (IVIS<sup>TM</sup> Perkin Elmer). For Post-treatment studies: mice were inoculated as described (n = 7/group), randomized and treated upon the detection of bioluminescent signal. Mice that showed tumor growth outside of the bone compartment were excluded from all analyses.

A detailed description of cell assays, histological and bone morphology analyses can be found in supplementary methods. Statistical analyses were performed with GraphPad Prism and all graphs display error bars that are SEM.

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#### **Author Contributions**

L.M.C. designed the study, performed the *in vitro* and *in vivo* studies and wrote the manuscript. A.A. developed the equations and implemented/analyzed *in silico* studies. M.M.B. optimized and contributed to *ex vivo* CT scan analysis. J.M.-S. contributed his clinical expertise on interpretation of the data. D.B. designed the study and wrote the manuscript. C.C.L. designed the *in vivo* components of the study, oversaw data collation and interpretation and wrote the manuscript.

#### **Additional Information**

Supplementary information accompanies this paper at http://www.nature.com/srep

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# Arturo Araujo, PhD

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# Profile

I'm currently in my last year of Post Doctoral training at the Integrated Mathematical Oncology (IMO) department at Moffitt Cancer Center, supervised by Dr. David Basanta (Cancer Evolution) and Dr. Conor Lynch (Tumor Biology). I model key aspects of cancer at the interphase between mathematics, biology and the clinic. My research deals with complex systems and emergent behaviors because I am interested in their potential to investigate complex biological ideas and counter intuitive results. Since joining the Moffitt Cancer Center in 2012, I have used all of my interdisciplinary training and viewpoints to successfully bridge the gap between theory, experiments and the clinic with integrated computational models in tight collaboration with the Lynch Lab and clinician Dr. Julio Pow-Sang. In my time here I have also strived to acquire the necessary skills to become a successful independent researcher. I'm fluent in English and Spanish and have experience in IT, programming, cell culture, teaching, art, graphic design, and leadership skills. Personal interests include complexity, evolutionary biology, technology, art, the advancement of humanity and films.

# Education

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**Conseil Européen pour la Recherche Nucléaire (CERN), Geneva, Switzerland** (2007) Teacher programme for the translation of science into classrooms

#### University of Notre Dame, Indiana, USA (2004-2005) One-Year Academic Exchange towards BSc in Physics

- Universidad de las Américas Puebla (UDLAP), Puebla, Mexico (2001-2006) Certificate in Art Studio
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- Araujo, A., Bentley, P. J. and Baum, B. (2010) Modelling the Role of Aneuploidy in Tumour Evolution. 12th International Conference on the Synthesis and Simulation of Living Systems. 2010. MIT Press.

# **Contributed Talks**

- Society for Mathematical Biology Annual Meeting 2016, Nottingham, UK. "Predictive modeling of treatment strategies for bone metastatic prostate cancer"
- The 11th American Institute of Mathematical Sciences Conference on Dynamical Systems, Differential Equations and Applications 2016, Orlando, FL, USA. "Predictive modeling of treatment strategies for bone metastatic prostate cancer"
- Society for Mathematical Biology Annual Meeting 2014, Osaka, Japan. "TGFβ in prostate to bone metastasis"
- Society for Mathematical Biology Annual Meeting 2013, Arizona, USA. "TGF $\beta$  inhibition in prostate to bone metastasis"
- Complex Systems Modelling and Simulation Workshop 2010, Odense, Denmark. "Modelling the Role of Chromosome Missegregation in Cancer Therapies"
- International Conference on the Synthesis and Simulation of Living Systems 2010, Odense, Denmark. "Modelling the Role of Aneuploidy in Tumour Evolution"

# **Invited Talks**

- Florida Prostate Cancer Research Symposium 2016, Clearwater, FL, USA Predictive modeling of treatment strategies for bone metastatic prostate cancer"
- Society for Mathematical Biology Annual Meeting 2015, Atlanta, GA, USA "Lessons From an Integrated Model of Prostate to Bone Metastasis"
- Florida Prostate Cancer Research Symposium 2015, Orlando, FL, USA " Lessons From an Integrated Model of Prostate to Bone Metastasis"
- Dartmouth Mathematics Departmental Colloquium 2015, Hanover, NH, USA "Lessons From an Integrated Model of Prostate to Bone Metastasis"
- Integrated Mathematical Oncology Seminar Series 2014, Tampa, FL, USA "How to build a cancer model".
- UDLAP Physics and Mathematics Annual Meeting 2014, Puebla, Mexico, "Improving treatment strategies for patients with metastatic castrate resistant prostate cancer through personalized computational modeling"
- UDLAP Physics and Mathematics Seminar 2013, Puebla, Mexico, "A Mathematical Framework Against Prostate Cancer
- IA Benemerita Universidad Autonoma de Puebla 2013, Puebla, Mexico, "Mathematics, Computer Science and the Advancement of Humanity"
- MRC-LMCB Laboratory for Molecular Cell Biology UCL Seminar Series 2013, London, UK. "Chromosome Missegregation in Cancer Development"
- Centre for Interdisciplinary Research in the Medical and Life Sciences Summer Seminar

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